

#### IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

Applicants: Craig A. Coburn et al.

Examiner: Yong Liang Chu

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For:

PHENYLCARBOXAMIDE BETA-SECRETASE INHIBITORS FOR THE TREATMENT OF ALZHEIMER'S DISEASE

Commissioner for Patents P. O. Box 1450 Alexandria, VA 22313-1450

#### DECLARATION OF MING-TAIN LAI UNDER 37 C.F.R. § 1.132

- I, Ming-Tain Lai, hereby declare as follows:
- 1. I am a citizen of the United States, and am over 21 years of age. A copy of my curriculum vitae is attached at Exhibit A.
- 2. In October 2002, HPLC assays of BACE1 (β-site amyloid precursor protein cleaving enzyme) were regularly conducted under my control and supervision at my laboratory at Merck's facility in West Point, Pennsylvania. Among the compounds tested were a series of phenylcarboxamide compounds, which were designed and synthesized by medicinal chemists working at Merck's West Point, Pennsylvania laboratories.
- 3. The BACE HPLC assay, which was a standard Merck assay, was developed by me and other biologists at Merck's West Point laboratories. The assay was designed to detect cleavage of a coumarin-labeled 10 mer peptide (coumarin-REVNFEVEFR), using either a Waters 2690 Alliance or Alliance HT HPLC instrument. The assay procedure is generally described in International application no. WO 2004/099376.
- 4. The BACE HPLC assays were conducted according to the following procedure. A reaction buffer was formed of the following ingredients:

MATERIAL ** * * * * * * * * * * * * * * * * *	AMOUNT (pl)
4X NaOAc, 200mM, pH 4.5	25
BSA, 1mg/ml (Bovine Fraction V, Sigma #9647)	2.0
EDTA, 150mM, pH 4.5	10
10% CHAPS(Pierce, #28300)	2.0
Deferoxamine Mesylate, 50mM (Sigma, #D9533)	2.0
b-BACE1 (20nM, 20mM Tris, pH 7.2)	10
H <sub>2</sub> O	31

- 5.  $8 \mu l$  of compound (in DMSO) was added to 90  $\mu l$  of the reaction buffer, and the resulting mixture was incubated at room temperature with shaking for 30 minutes.
- 6. Thereafter, 2  $\mu$ l of the substrate coumarin-CO-REVNFEVEFR (50 $\mu$ M) (as described in WO 2004/099376) was added to the mixture. The resulting mixture was maintained at 25°C for 30 minutes with shaking. The reaction was quenched with 25 $\mu$ l of 1M Tris-HCl pH8.0.
- 7. Samples of the mixture were then centrifuged in a tabletop centrifuge at 15K. For analysis with the Alliance HPLC instrument, 60  $\mu$ l of the supernatant was removed. For analysis with the Alliance HT instrument, 50  $\mu$ l of the sample was passed through a filtration system (Millipore, 0.22  $\mu$ m hydrophilic) prior to HPLC analysis.
- 8. The HPLC conditions involved an Xterra RP18 column (3.5  $\mu$ m, 2.1 x 150 mm). The mobile phase consisted of solvent A (0.05% trifluoro acetic acid in water) and solvent B (0.045% trifluoracetic acid in acetonitrile), according to the following gradient:

Time (minutes)	Percent Solvent B
0	19
3	25
4	95
5	19

Sample injection volumes were 50  $\mu$ L for the Alliance and 25  $\mu$ L for the Alliance HT. Detection was measured at 340 nm (excitation) and 440 nm (emission). Percent inhibition was measured according to the following formula:

(1-(area of product peak) of (E+S+compound)/area of product peak of (E+S)) x 100

9. The results of the HPLC assay for selected phenylcarboxamide compounds are set forth below:

COMPOUND	Date of Testing	Inhibition of BACE1 (nM)
0,0 H <sub>3</sub> C <sub>N</sub> -S <sup>2</sup> CH	June 7, 2002	3
F OH		
à, ö "		
F F 0-		
F <sup>2</sup> Yo <sup>-</sup>		
0		
4,0 4,0 4,0	June 7, 2002	220
FY OH		
. 11		
}	1	

H,C., N,S, CH,  O,O,O  H,C., N,S, CH,  OH  OH  OH  OH  OH  OH	June 10. 2002	1
H,C, N, S, CH, OH, N, O	July 9, 2002	3.5
H <sub>2</sub> C S N CH <sub>3</sub> OH  OF F	July 17, 2002	46
H <sub>2</sub> C-S N-CH <sub>3</sub> OH  OH  OH  OH  OH  OH  OH  OH  OH  O	October 7, 2002	11

10. I further declare that all statements made herein of my own knowledge are true, and that all statements made on information and belief are believed to be true; and further that these statements are made with knowledge that willful false statements and the like so made are punishable by fine or imprisonment or both, under § 1001 of Title 18 of the United States Code, and that such willful false statements may jeopardize the validity of the instant application or any patent issued thereon.

Ming-Tain Lai

Dated: May 8, 2006



#### **CURRICULUM VITAE OF MING-TAIN LAI**

#### **PERSONAL**

A. Name: Ming-Tain Lai

B. Home Address: 52 Douglass Road

Lansdale, PA 19446

### II. <u>EDUCATION</u>:

School	Dates	Major	Degree
Tunghai University Taiwan	1977-1981	Chemistry	B.S.
National Taiwan Normal University, Taiwan	1981-1983	Analytical Chemistry	M.S.
University of Minnesota	1987-1992	Bioorganic Chemistry	Ph.D.

#### III. MRL EMPLOYMENT HISTORY

<u>Title</u> <u>From</u> - <u>To</u>

Research Fellow 8/30/01 - present

Senior Research Biochemist 8/30/95 - 8/30/01

#### IV. NON-MERCK EMPLOYMENT HISTORY

Postdoctoral Research Associate, 1992-1995 Massachusetts Institute of Technology Supervisor: Professor JoAnne Stubbe

## V. <u>SOCIETY MEMBERSHIPS</u>

American Chemical Society

#### VI. PUBLICATIONS IN PEER REVIEWED JOURNALS

- 1. Lai, M-t.; Shih, J-S., "Mercury (II) and Silver (I) Ion-Selective Electrodes Based on Dithia Crown Ether," Analyst 1986, 111, 891-895.
- 2. Lai, M-t.; Lin, W-M.; Chu, Y-H.; Chen, Y. S-l.; Kong, K-S.; Chen, C-w., "The Mechanism of Color Reversion in Soybean Salad Oil," J. Am. Oil Chem. Soc. 1989, 66, 565-571.
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#### VIII PATENTS

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- 2. Hazuda, D.; Dodson, E. C; Lai, M.-t; Xu, M.; Shi, X.-P.; Simon, A, J.; Wu, G.; Li, Y.; Register, R. B.,. Assays Using Amyloid Precursor Proteins with Modified Beta-Secretase Cleavage Sites to Monitor Beta-Secretase Activity. Filed in February, 2003. Al Published: 20031023 as US20030200555 A1
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- 4. Crouthamel, M. C.; Gardell, S. J.; Huang, Q.; Lai, M.-t.; Li, Y., Gamma-3 protease, Application No. PCT/US02/26969, Filed Aug. 8, 2002, Publication No. WO 03/018050 A1

#### IX. <u>ABSTRACTS</u>

- 1. Lai, M-t., Oh, E., Liu, L-d., Li, D., Liu, H-w., "Mechanistic Study on the Inactivation of General Acyl-CoA Dehydrogenase by a Metabolite of Hypoglycin A," XI Midwest Enzyme Chemistry Conference, University of Illinois, Chicago, IL, 1989.
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- 3. Lai, M-t., Oh, E., Liu, L-d., Li, D., Liu, H-w., "Mechanistic Study on the Inactivation of General Acyl-CoA Dehydrogenase by a Metabolite of Hypoglycin A" XI Midwest Enzyme Chemistry Conference, University of Illinois, Chicago, IL, 1991.
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- 5. Li, D., Oh, E., Lai, M-t., Zhou, H-1., Becker, D.F., Stankovich, M.T., Liu, H-w., "Studies of the Inactivation of Short-Chain Acyl-CoA Dehydrogenase by Drivatives

- of Methylenecyclopropaneacetyl-CoA" XIII Midwest Enzyme Chemistry Conference, Loyola University Chicago, ILL, 1993.
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- 7. Brady, S; Bruce, J.; Singh, S.; Crouthamel, M.-C.; Holloway, K. M.; Coburn, C.; Vacca, J. P.; Shafer, J.; Hazuda, D. "Development of BACE 1 Inhibitors" 9<sup>th</sup> International Conference on Alzheimer's Disease and related Disorders, Philadelphia, PA, July 17-22, 2004

# X. INVITED LECTURES

3/11/9	of the Ina	nt of Chemistry, National Chung-Cneng University, "Mechanistic Study stivation of Medium Chain Acyl-CoA Dehydrohegenase by ecyclopropane)acetyl-Co-A"
3/15/9	the Inactiv	nt of Chemistry, National Chiao-Tung University, "Mechanistic Study of ration of Medium Chain Acyl-CoA Dehydrogenase by ecyclopropane)acetyl-Co-A"
3/18/9	the Inactiv	nt of Chemistry, National Tsing-Hua University, "Mechanistic Study of ration of Medium Chain Acyl-CoA Dehydrogenase by ecyclopropane)acetyl-Co-A"
3/22/9	Inactivation	nt of Chemistry, National Taiwan University, "Mechanistic Study of the on of Medium Chain Acyl-CoA Dehydrogenase by ecyclopropane)acetyl-Co-A"
2/13/9	Stable, No	nt of Chemistry, National Taiwan University, "Characterization of a vel Norcaradiene Adduct Resulting from the Inactivation of Thymine see by 5-Ethynyluracil"
2/16/9	a Stable, 1	nt of Life Science, National Tsing-Hua University, "Characterization of Novel Norcaradiene Adduct Resulting from the Inactivation of Thymine see by 5-Ethynyluracil"
2/22/9	of a Stabl	nt of Chemistry, National Taiwan Normal University, "Characterization e, Novel Norcaradiene Adduct Resulting from the Inactivation of Hydroxylase by 5-Ethynyluracil"
12/8/2		Cargets and Drug Discovery Conference, Strategic Enstitute, "Development of BACE 1 Inhibitors"
7/21/		ase, 9 <sup>th</sup> International Conference on Alzheimer's Disease and Related "Development of BACE 1 Inhibitors"